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(54) SOLID POLYMER ELECTROLYTE

(57) An object of the present invention is to provide a practical solid polymer electrylet that exhibits excellent thermal characteristics, physical characteristics, and ionic conductivity, and in particular, to provide a completely solid electroyle and a copolymer composition that functions as the substrate for producing such an electroylet, a solid polymer electroylet comprising an electroylet as all, and a copolymer in which a block chain A containing a repeating unit represented by a formula (i) shown below.

a block chain B containing a repeating unit represented by a formula (II) shown below:

$$\begin{array}{c|c}
R_6 & R_8 \\
\hline
C & C \\
R_7 & R_9
\end{array}$$

(wherein, R_g represents an aryl group), and a block chain C containing a repeating unit represented by a formula (III) shown below:

$$\begin{array}{c|cccc}
 & R_{10} & R_{12} \\
\hline
 & C & C \\
R_{11} & R_{13} & & & & & & & & & \\
\end{array}$$

(wherein, R₁₃ represents an aryl group or a heteroaryl group) are arranged in the sequence B, A, C.

EP 1 553 117 A1

Description

TECHNICAL FIELD

5 [0001] The present invention relates to a solid polymer electrolyte that is particularly useful for use within secondary batteries.

BACKGROUND ART

- (0002] Although batteries have long been used. They have recently been recognized as one of the extremely important base components, together with semiconductors and liquid crystal display elements, in the information age. Practically in the cases of portable telephones and notabook type personal computers, the demands for improved battery performance and reduced battery weight are strong, and lithium ion batteries are attracting considerable attention as one type of battery capable of meeting these demands. Lithium ion batteries are destribed another series of the strength of the provision of the destribed provision for the future. (2003) In electrochemical elements such as nickel-cannium batteries, and offer considerable promise for the future reliability in contrast, solid electrolytes as primary batteries, secondary batteries and capacitors, liquids have reliability in contrast, solid electrolytes do not suffer from these drawbacks, and the application of such solid electrolytes reliability in contrast, solid electrolytes do not suffer from these drawbacks, and the application of such solid electrolytes to a variety of electrochemical elements not only simplifies the production of the element, but also enables reductions are in the size and weight of the element, and because of the lack of any leakage problems, enables the provision of a highly reliable element.
 - [0004] Accordingly, in the field of lithium on battery research, the research and development of new solid electrolytes, and in particular the research and development of solid polymer electrolytes that are lightweight, flexible and readily orccessed, is being actively oursued.
- 25 (0005) Almost all polymer compounds are insulators, but since the announcement that certain polymer materials such as polyterylene oxide (PEQ) are capable of forming a crystalline complex with electrohyte salts such as inhum salts, resulting in a high ionic conductivity much attention has been focused on research into solid polymer electrohytes that use PEQ or other polyativityone oxide, or polyethyrenimics or polytesphagenes that contain entitler in dissociation groups as the electrohyte matrix. There have been many publications of research relating to solid polymer electrohytes in which a polyatilysiene oxide represents one component of the matrix, and the lonic conductivity at room cities of the properties of the properties of the properties of the properties of the matrix must be increased, but unfortunately this leads to a marked deterioration in the strength and heat resistance of the electrohyte film, meaning producing a practical solid electrohyte has been difficult. Moreover, the ionic conductivity also decreases dramatically if the temperature falls to 0°C or lower (see Jepanese Unexamined Patent Angloadents. Pixel Publication No. Heis 1-20912, and J. Appl. Electroherm. No. 5, 63 to 96 119730.
- [0006] A solid polymer electrolyte in which the matrix substrate comprises an ABA type triblock copolymer produced by oppolymerzing methoxypolyethylene glycol monomethacylate (A) and styrene (B) via a living anion-c polymerization has been proposed as a subtable solid polymer electrolytic (see Makromol, Chem, 190, 1058 to 1078 [1389]).
- [0007] However, the homopolymer of the methoxypolyethylene glycol monomethacrylate of the component (A) is a liquid at room temporature, even at very high molecular weights, meaning that in order to ensure that the ABA copolymer forms a solid electrolyte matrix substrate, the quantity of the component (A) must be restricted. However, this means that the shape and size of the PEO domain, which functions as the diffusion and transport space for the lithium ions, is also restricted, and as a result, the ionic conductivity at 40°C was a somewhat unsatisfactory (3° SC/m.

45 DISCLOSURE OF INVENTION

- [0008] An object of the present invention is to provide a solid polymer electrolyte that utilizes a microphase separated structure to provide excellent ionic conductivity, thermal characteristics, and physical characteristics.
- [0009] As a result of intensive investigations aimed at resolving the problems described above, the inventors of the present invention discovered that in a triblock copolymer containing a block chain formed from alloxypolyethylene glycol monotimeth)acrylate units, if a rigid block chain such as polyetyrene is provided at either one, or both terminals of the alkoxypolyethylene gycol mono(meth)acrylate chain, then the pseudo-cross-linking effect provided by the rigid block chain enables sclidification to be achieved even when the quantity of alkoxypolyethylene glycol mono(meth) acrylate units is increased, thus enabling a higher ionic conductivity to be achieved, and enabling the inventors to conclude the present inventions.

[0010] In other word, a first aspect of the present invention relates to the following:

(1) A solid polymer electrolyte comprising an electrolyte salt, and a copolymer in which a block chain A containing

a repeating unit represented by a formula (I) shown below:

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(wherein R_1 to R_2 each represent, independently, a hydrogen atom or a hydrocarbon group of C1 to C10, R_1 and R_2 may be bonded together to form a ring. R_{4a} and R_{4b} each represent, independently, a hydrogen atom or a methyl group. R_2 represents a hydrogen atom or a methyl group, R_2 represents a hydrogen atom, a hydrocarbon group, an acyl group, or a silyl group, in represents an integer from 2 to 100, and individual R_{4a} and R_{4b} groups may be either the same or different), a block chain B containing a repeating unit represented by a formula (fill) shown below:

(wherein B_6 to B_6 each represent, independently, a hydrogen atom or a hydrocarbon group of C1 to C10, and B_6 represents an anyl group), and a block chain C are arranged in the sequence B_1A_1 .

The present invention also relates to the following aspects:

(2) A solid polymer electrolyte according to the first aspect, wherein the block chain C contains a repeating unit represented by a formula (iii) shown below:

$$\left(\begin{array}{c|c} R_{10} & R_{12} \\ \hline C & C \\ R_{11} & R_{13} \end{array} \right) \quad \ldots \quad (\text{III})$$

(wherein R_{10} to R_{12} each represent, independently, a hydrogen atom or a hydrocarbon group of C1 to C10, and R_{13} represents an anyl group or a heteroaryl group).

(3) A solid polymer electrolyte according to either one of the first and second aspects, wherein the block chains A to C form a copolymer with a B-A-C bonding sequence.

(4) A solid polymer electrolyte according to any one of the first through third aspects, wherein the degree of polymerization of the repeating unit represented by the formula (f) is at least 10.

(5) A solid polymer electrolyte according to any one of the first through fourth aspects, wherein the degree of polymerization of the repeating unit represented by the formula (II) is at least 5.

(6) A solid polymer electrolyte according to any one of the second through fifth aspects, wherein the degree of polymerization of the repeating unit represented by the formula (III) is at least 5.

(7) A solid polymer electrolyte according to any one of the first through sixth aspects, wherein the value of m in the formula (I) is an integer from 5 to 100.

(8) A solid polymer electrolyte according to any one of the first through seventh aspects, wherein the value of m in the formula (I) is an integer from 10 to 100.

(9) A solid polymer electrolyte according to any one of the second through eighth aspects, wherein the group R₁₃

in the formula (III) is an anyl group, and the degree of polymerization of the repeating unit represented by the formula (III) is at least 5.

(10) A solid polymer electrolyte according to any one of the first through ninth aspects, wherein the molar ratio ((i) ((il)-(i)) obsvere the exposaling units represented by the formula (i), and the combined total of the repeating units represented by the formula (ii) and the repeating units within the block chain C is within a range from 1/30 to 30/1. (11) A solid polymer delectrolyte according to any one of the second through ninth aspects, wherein the molar ratio ((ii)((iii)-(iii))) between the repeating units represented by the formula (i), and the combined total of the repeating units represented by the formula (iii) is within a range from 1/30 to 30/1.

(12) A solid polymer electrolyte according to any one of the first through eleventh aspects, wherein the number average molecular weight of the copolymer is within a range from 5,000 to 1,000,000.

(13) A solid polymer electrolyte according to any one of the first through twelfth aspects, which exhibits a microphase separated structure.

[14] A solid polymer electrolyte according to any one of the first through thirteenth aspects, wherein the electrolyte salt is one or more materials selected from a group consisting of alkali metal salts, quaternary ammonium salts, quaternary prospondinum salts, sustrain metal salts, and protonic acids.

(15) A solid polymer electrolyte according to any one of the first through thirteenth aspects, wherein the electrolyte salt is a lithium salt.

The present invention also relates to a sixteenth aspect described below:

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(16) A copolymer in which a block chain A containing a repeating unit represented by a formula (I) shown below:

(wherein R_1 to R_2 each represent, independently, a hydrogen atom or a hydrocarbon group of C1 to C10, R_1 and R_2 , and y be bonded together to form a ring, R_4 and R_4 , each represent, independently, a hydrogen atom or a methyl group, R_3 represents a hydrogen atom or a methyl group, R_3 represents a hydrogen atom or a methyl group, an acyl group, or a siyl group, or represents an integer from 2 to 100, and individual R_4 , and R_6 , groups may be either the same or different), a block chain B containing a repeating unit represented by a formula (II) shown below:

(wherein R_6 to R_6 each represent, independently, a hydrogen atom or a hydrocarbon group of C1 to C10, and R_9 represents an anyl group), and a block chain C containing a repeating unit represented by a formula (III) shown below:

$$\begin{array}{c|c}
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 &$$

(wherein B₁₀ to B₁₂ each represent, independently, a hydrogen atom or a hydrocarbon group of C1 to C10, and B₁₃ represents an anyl group or a heteroaryl group) are arranged in the sequence B, A, C.

The present invention also relates to the following aspects:

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- (17) A copolymer according to the sixteenth aspect, wherein the block chains A to C are bonded together in a S-A-C sequence.
- (18) A copolymer according to either one of the sixteenth and seventeenth aspects, wherein the degree of polymerization of the repeating unit represented by the formula (i) is at least 10.
- (19) A copolymer according to any one of the sixteenth through eighteenth aspects, wherein the degree of polymerization of the repeating unit represented by the formula (th) is at least 5.
- (20) A copolymer according to any one of the sixteenth through nineteenth aspects, wherein the degree of polymerization of the repeating unit represented by the formula (III) is at least 5.
 - (21) A copolymer according to any one of the sixteenth through twentieth aspects, wherein the value of m in the formula (1) is an integer from 5 to 100.
 - (22) A copolymer according to any one of the sixteenth through twentieth aspects, wherein the value of m in the formula (I) is an integer from 10 to 100.
 - (23) A copolymer according to any one of the sixteenth through twenty second aspects, wherein the group R₁₃ in the formula (III) is an anyl group, and the degree of polymerization of the repeating unit represented by the formula (III) is at least 5
- (24) A copolymer according to any one of the sixteenth through twenty third aspects, wherein the motar ratio ((I)/(III)+(III)) between the repeating units represented by the formula (I), and the combined total of the repeating units represented by the formula (III) is within a range from 1/30 to 30/f.
 - (25) A copolymer according to any one of the sixteenth through twenty fourth aspects, wherein the number average molecular weight of the copolymer is within a range from 5,000 to 1,000,000.
- 25 (26) A copolymer according to any one of the sixteenth through twenty fifth aspects, which exhibits a microphase separated structure.
 - The present invention also relates to the following aspects.
 - (27) A method of producing a copolymer according to any one of the sixteenth through twenty sixth aspects, which utilizes a living radical polymerization in which a transition metal complex is used as a catalyst, and an organichal-open compound comprising 1 or more halogen atoms is used as an initiator.
 - (28) An ion conductive film containing a polymer comprising a polymer segment (P1) that displays ionic conductivity and a polymer segment (P2) that displays no ionic conductivity, wherein the film exhibits a network-type microphase separated structure.
 - (29) An ion conductive film according to the twenty eighth aspect, wherein the polymer comprising P1 and P2 is a polymer in which the segments are arranged in a P2, P1, P2 sequence.
 - 1) Copolymer used in the Solid Polymer Electrolyte of the Present Invention

lecular weight linkage group such as an oxygen atom or an alkylene group.

- [0011] A copolymer used in a solid polymer electrolyte of the present invention is a copolymer comprising a block of chain A containing a repeating unit represented by the formula (f), a block chain B containing a repeating unit represented by the formula (fl), and an arbitrary block chain C, wherein these block chains are arranged in the sequence B, A, C, in this case, the block chain B and the block chain C may be either the same or different.
 - [0012] The statements that the block chains A and B contain repeating units represented by the formulas (i) and (ii) respectively include those cases in which the block chains contain other repeating units as structural units in addition to the identified repeating units, as well as those cases in which no other structural units exist within the chains. Furthermore, in those cases where a block chain also contains other repeating units as structural units, there are no particular restrictions on the form of the polymer generated with the repeating unit represented by the above formula, and random polymers, block polymers and alternating polymers are all possible.
- [0013]. Furthermore, the statement that the block chains are arranged in the sequence B, A. C. includes those cases in which the blocks are bonded together directly, as well as those cases in which the blocks are bonded together via other structural units such as linkage groups or other polymer chains. In those cases where a copolymer also comprises another polymer chain as a structural unit, this chain may be either a homopolymer, or a binary or higher order copolymer, are in the case of a copolymer, there are no particular restrictions on the bonding within the copolymer and random, block, and gradient copolymers in which the component ratio gradually changes, are all suitable. Of the cases cases/field above, those cases in which each block chains is bonded in this sequence B-A.C are preferred. In such cases, the term "bonded" means that the block chains are oither bonded logether excelly, or bonded together via a low.
 - [0014] In the repeating unit represented by the formula (I), the groups R₄ to R₅ each represent, independently, a

hydrogen atom or a nydrocarbon group of 1 to 10 carbon atoms such as a methyl group, ethyl group, n-propyl group, isopropyl group, n-butyl group, sec-butyl group, isobutyl group, b-butyl group, paphthyl group, naphthyl group, or benzyl group. Eurthermore, R, and R, mety be bonded together to form a ring.

[0015] The groups R_{4a} and R_{4b} each represent, independently, a hydrogen atom or a methyl group, and m represents an integer from 2 to 100, and preferably an integer from 5 to 100, and even more preferably an integer from 10 to 100. The values of m within each repeating unit may be either the same or different, and individual R_{4a} groups and R_{4b} groups may be either the same or different.

[0016] R₅ represents a hydrogen atom; a hydrocarbon group such as a methyl group, ethyl group, n-propyl group, isopropyl group, n-busyl group, see-busyl group, isobusyl group, is-busyl group, n-hexyl group, phenyl group, substituted phenyl group, on naphthyl group; an acyl group such as a formyl group, acetyl group, propionyl group, busyryl group, or benzyll group; or a silyl group such as a trimethylistyl group or at butyldimethylsilyl group.

[0017] The groups R₁ to R₂ may contain a substituent group at a suitable carbon atom, and specific examples of this substituent group include a halogen atom such as a fluorine atom, chlorine atom, or bromine atom, a hydrocerbon group such as a mothyl group, othyl group, n-propyl group, phonyl group, naphthyl group, or benzyl group, an acyl group such as an acetyl group or a benzely group, a hydrocerbon oxy group such as a nitrite group, nitro group methoxy group, or phenoxy group, as well as other groups such as a methylthio group, methylsulfinyl group, methylsulfonyl group, amino group, of methylsulfonyl group, dimiditylamino group, or anilino group.

[0018] The degree of polymerization of the repeating unit represented by the formula (f) varies depending on the value of m, although values of at least 10 are preferred, and values of 20 or greater are even more preferred. Specific examples of the repeating unit represented by the formula (f) include the compounds listed below. The compounds listed represent monomers that can be considered as giving rise to the repeating unit represented by the formula (f). Furthermore, these repeating units can be employed either singularly, or in mixtures of two or more different repeating units.

[0019] 2-methoxyethyl (meth)acrylate, 2-ethoxyethyl (meth)acrylate, 2-methoxypropyl (meth)acrylate, 2-ethoxypropyl (meth)acrylate, methoxypopylethylene glyocl (meth)acrylate, (in which the number of ethylene glyocl of meth)acrylate, phenoxypolyethylene glyocl (meth)acrylate, phenoxypolyethylene glyocl (meth)acrylate, phenoxypolyethylene glyocl (meth)acrylate, phenoxypolyeropylene glyocl (meth)acrylate, phenoxypolyeropylene glyocl (meth)acrylate, phenoxypolyeropylene glyocl (meth)acrylate, polyethylene glyocl mono(meth)acrylate, polyethylene glyocl mono(meth)acrylate, polyethylene glyocl mono(meth)acrylate, polyethylene glyocl mono(meth)acrylate, glyocl mono(meth)acrylate, glyocl mono(meth)acrylate, glyocl mono(meth)acrylate, stearoxypolyethylene glyocl mono(meth)acrylate, the Blemer PME series of compounds (monomers of the formula (i) in which R₁ = R₂ = a Hydrogen atom, R₂ = a methyl group, and m = 2 to 90) (manufactured by NIPPON ICL. & PAT Corporation), acrylocypolythylene glyocl (meth)acrylate, betrayolyoxypolyethylene glyocl (meth)acrylate, butlylidenthylidene glyocl (meth)acrylate, butlylidenthylidene glyocl (meth)acrylate, butlylidenthylidene glyocl (meth)acrylate, butlylidenthylidene glyocl (meth)acrylate, methylishylicyypolyethylene glyocl (meth)acrylate, methylishylicyypolyethylene glyocl (meth)acrylate, methylishylidene glyocl

polyethylene glycol cyclohexene-1-carboxylate, and methoxypolyethylene glycol cinnamate. [0020] In the respeating unit represented by the formula (II), the groups R_g to R_g each represent, independently, a hydrogen atom or a hydrocarbon group of 1 to 10 carbon atoms such as a methyl group, ethyl group, n-propyl group, isopropyl group, n-butyl group, etholyl group, etholyl group, phenyl group, nabhthyl group, or benzyl group, and the group R_g represents an anyl group such as a phenyl group, substituted phenyl group, naphthyl group, or antiracenty group.

[0021] Furthermore, the groups R₈ to R₈ may contain a substituent group at a suitable carbon atom, and specific examples of this substituent group natiotide a halogen atom such as a fluorine atom, or hiorine atom, or bromline atom, or hiorine atom, or hidden, and hidden at his description, and an atom at the suitable at a nativity group, or a suffice group, nativity group, or phenoxy group, as well as other groups such as a methythilo group, methylsidinyl group, methylsidinyl group, armine group, dimethylamine group, or aniline group.

[0022] The degree of polymerization of the repeating unit represented by the formula (II) is preferably at least 5, and even more preferably 10 or greater. Specific examples of the repeating unit represented by the formula (II) include the compounds listed below. The compounds listed represent monomers that can be considered as giving rise to the repeating unit represented by the formula (II). Furthermore, these repeating units can be employed either singularly, or in mixtures of two or more different repeating units.

[0023] Aryl compounds such as styrene, o-methylstyrene, p-methylstyrene, p-t-butylstyrene, c-methylstyrene, p-t-butoxystyrene, m-t-butoxystyrene, a-dimethylstyrene, m-t-birostyrene, p-t-birostyrene, 4-carboxystyrene, vinylarisloe, vinylaris

55 [0024] The block chain C is a block chain containing an arbitrary repeating unit, although examples of preferred block chains include those containing a repeating unit represented by the formula (III).

[0025] In the repeating unit represented by the formula (III), the groups R₁₀ to R₁₂ each represent, independently, a hydrogen atom or a hydrocarbon group of 1 to 10 carbon atoms such as a methyl group, elhyl group, n-propyl group,

isopropyl group, n-butyl group, sec-butyl group, isobityll group. E-butyl group, phenyl group, nachthyl group, or benzyl group. The group R₁₃ represents en anyl group such as a phenyl group, substituted phenyl group, nachthyl group, or antifracenyl group, or a heteroaryl group such as a 2-pyridyl group or 4-pyridyl group.

[0026] Furthermore, the groups R₁₀ to R₁₅ may contain a substituent group at a suitable carbon atom, and specific examples of this substituent group include a halogen atom such as a fluorine atom, or bromine atom, a hydrocarbon group such as a mathyl group, or benzyl group, phenyl group, naphthyl group, or benzyl group, an acyl group such as an seetly group or a benzolf group, an dayl group such as an atom at a substitution of group, mathy group, or phenoxy group, as evel as other groups such as a mathylthic group, methylsulfnyl group, methylsulfnyl group, methylsulfnyl group, amino group, dimethylsming group, or antino group.

[0027] The degree of polymerization of the repeating unit represented by the formula (III) is preferably at least 5, and even more preferably 10 or genelaric Specific examples of the repeating unit represented by the formula (III) include the compounds lated below. The compounds lated represent monomers that can be considered as giving rise to the repeating unit represented by the formula (III). Furthermore, these repeating units represented by the formula (III) can be employed other singularity or it mixtures of two or more different repeating units.

[0028] Aryl compounds such as styrene, o-methylstyrene, p-methylstyrene, p-t-butylstyrene, α-methylstyrene, p-t-butystyrene, α-methylstyrene, p-t-butystyrene, α-methylstyrene, α-methylstyre

[0029] A copolymer of the present invention may also contain repeating units that are different from the repeating units represented by the formulas (i) to (iii) as structural units, and examples of these other repeating units include the compounds listed below. Furthermore, these repeating units can also be used within the block chain. In those cases where the block chain. I does not contain a repeating unit represented by the formula (iii). The compounds listed represent monomers that can be considered as giving rise to the repeating units. These repeating units can be employed either singularly, or in mixtures of two or more different repeating units.

[0030] Suitable compounds include (meth)acrylates such as methyl (meth)acrylate, ethyl (meth)acrylate, bottyl (meth)acrylate, bottyl (meth)acrylate, bottyl (meth)acrylate, bottyl (meth)acrylate, churyl (meth)acrylate, methyl (methylacrylate, methyl (meth)acrylate, methyl (methylacrylate, m

[0031] In addition, repeating units that contain an intramolecular double bond capable of undergoing copolymerization with a monomer that generates a repeating unit represented by one of the formulae (1) to (1), and also contain at least one functional group selected from a group consisting of a hydroxyl group, carboxyl group, epoxy group, acid anhydride group and amino group may also be incorporated as structural units, and specific examples of such repeating units include the compounds listed below. The compounds listed represent monomers that can be considered as giving rise to the equivalent repeating unit. These repeating units can be employed either singularly, or in mixtures of two or more different repeating unit.

[0032] The molar ratio (II)(III)+CI) between the repeating units represented by the formula (I), and the combined total of the repeating units represented by the formula (II) and the repeating units within the block chain C is preferably within a range from 1/30 to 30/1. If the preportion of the repeating units represented by the formula (I) is less than 1/30, then a satisfactory level of conductivity cannot be achieved, whereas if the proportion exceeds 30/1, then satisfactory thermal characteristics annot be achieved. If the proportion represented by the formula (II) and the repeating units of the block chain C is less than 1/30, then satisfactory thermal characteristics and physical characteristics cannot be achieved whereas if the proportion exceeds 30/1, then a satisfactory level of conductivity cannot be achieved. Furthermore, in those cases where the block chain C contains a repeating unit represented by the formula (III), these repeating units represented by the formula (III) as a mirry by a insert by a insert of within the above ratio in place of the repeating units represented by the formula (III) as mirry by a insert of within the above ratio in place of the repeating units from the block chain C because where the solution of the proposition of the propos

[0033] Although there are no particular restrictions on the number average molecular weight of a copolymer of the present invention, values within a range from 5,000 to 1,000,000 are preferred. If the number average molecular weight is less than 5,000, then the thermal characteristics and physical characteristics tend to deteriorate, whereas if the value exceeds 1,000,000, the moletability and film forming characteristics tend to deteriorate. Furthermore, there are no particular restrictions on the ratio (MwM/In) between the weight average molecular weight (Min), although in order to ensure formation of the microphase separated structure described below, the ratio is proferebly within a range from 1,01 to 2,50, and even more referrably from 1,01 to 1,50.

[0034] There are no particular restrictions on the structure of the copolymer of the present invention, provided the block chans A, B, and C are arranged in a specific sequence, although in order to maintain a high level of ionic conductivity when used as a solid polymer electrolyte, the film structure preferably exhibits a microphase separated structure, and even more preferably a network type microphase separated structure.

2) Method of Producing the Copolymer

[0035] In those cases where the copolymer of the present invention comprises repeating units of the formulas (IV) and (V) shown below, as well as a repeating unit represented by the general formula (III) as the block chair C, the copolymer can be produced from a compound represented by the formula (VI) shown below, using known methods such as a living radical polymerization that uses a transition metal complex as the catalyst and an organchalegon compound comprising one or more hadegen atoms as the polymerization interface, a living radical polymerization using a stable radical, or a living anionic polymerization. Of these, a living radical polymerization using a transition metal complex as the catalyst and an organohalogen compound comprising one or more hategen atoms as the polymerization initiator is preferred.

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[0036] The groups B_4 to B_{13} in the formulas (IV) to (VI) are as defined above. A more specific description of the methods reveals the following options:

- (i) a method in which a compound represented by the formula (IV) is reacted via a living radical polymerization method using a bifunctional infiliator, and the thus obtained macroinitiator comprising bifunctional block chains and the like is then further reacted with a monomer that generates another block chain, thus sequentially extending the block chain.
- (ii) a method in which, with the exceptions of using a compound represented by the formula (V) instead of the compound represented by the formula (IV), and using a monofunctional artillator, reaction is conducted in the same manner as ii). Thus sequentiatly extending the block chair from the terminal, and
- (iii) a method in which each block chain, or a portion of each block chain, is polymerized using a predetermined sequence, and the block chains are subsequently subjected to a coupling reaction.
- 38 [0037] A living radical polymerization can be conducted using a transition metal complex as the catalyst and an organishatogen compound comprising one or more halogen atoms within each motecule as the polymerization initiator. [0038] The central metal within the transition metal complex is preferably manganase, thenium. Iron, ruthenium, rhodium, nickel, or an element from group? To 11 on the periodic table such as copper (using the periodic table disclosed in "Handbook of Chemistry, I Basic. 4th revised odition" (1983), odited by The Chemical Society of Japan; Of these metals, ruthenium is particularly preferred. There are no particular restrictions on the ligands that can be coordinated with the metal to form the metal complex, although examples of preferred ignaris include triary phosphines of 18 to 5 carbon atoms such as triphenyphosphine and frinaphthylphosphine, triarlyphosphines of 3 to 16 carbon atoms use as triphenyphosphine and trinaphthylphosphine such as triphenyphosphine, diphonyphosphine, balogen atoms such as isodine, bromine, and chlorine, as well as carbon monoxide, hydrogen atoms, cycloperiadeine, cyclope
 - including alkyl groups (including C1 to C4 alkyl groups such as methyl groups and ethyl groups), alkenyl groups (including C2 to C5 alkenyl groups such as vinyl groups and ethyl groups), alkenyl groups, alk

[0039] Of the ligands listed above, the hydrocarbon ligands may also comprise any of a variety of substituents,

[0040] Furthermore, in addition to the ligands described above, the transition metal complex may also contain a

hydroxyl group, an alkoxy group (such as a methoxy, othoxy, propoxy, or butoxy group), an acyl group (such as a nearly) or projetory group), an akovyaethoxyl group (such as a methoxycathory) or ethoxycathoryl group), all ediketone group such as eccytacetonate, a B-ketoester group such as acetylacetate, a pseudo-hatogen group [such as a CN, thioxyanate (SCN), selencocyanate (SeCN), tellurocyanate (TeCN), SCSN₃, OCN, ONC, or azide (N₃) group), an oxygen atom. H₂O, or a nitrogen-containing compound [such as NH₃, NO, NO₂, NO₃, ethylenodiarnine, dishylenotriarnine, trbutylarnine, 1,3-disporpolyl-4,5-dimethyllmidazed-2-yldine, pyridine, phenanthroline, or substituted phenanthroline, 22-8°,2'-4'repyridine, pyridinine, cross-sinked aliphated dismine, 4-4'-dif-chonyl-2'-bibyyridine, byyridine with coordinated thiocyanate, O, S, Se or Te, altylimineopyridine, altylybyridinylarnine, alkyl substituted tripyridine, difakt/arninolakt/ovarine, ethylenediarnineodovridine or trifovridin/repetrivilarnine).

[0041] Using transition metal complexes with ruthenium as the central metal as representative examples, specific examples of the transition metal complex used in producing a copolymer of the present invention include dichlorotris (triphenylphosphine) ruthenium, dichlorotris(tributylphosphine) ruthenium, dichloro(trialkylphosphine)p-cymene ruthenium, dichloro-diffricymenephosphine)styryl ruthenium, dichloro(cyclooctadlene) ruthenium, dichlorobenzene ruthenium, dichtoro-p-cymene ruthenium, dichtoro(norbornadiene) ruthenium, cis-dichtorobis(2,2'-bipyridine) ruthenium, dichlorotris(1,10-phenanthroline) ruthenium, carbonylchlorohydridotris(triphenylphosphine) ruthenium, chlorocyclopentadienylbis(triphenylphosphine) ruthenium, chloroindenylbis(triphenylphosphine) ruthenium, and dihydrotetrakis(triphenylphosphine) ruthenium, and of these, dichlorotris(triphenylphosphine) ruthenium, chloroindenylbis(tripherrylphosphine) ruthenium, and dihydrotetrakis(triphenylphosphine) ruthenium are preferred. In addition, specific examples of other transition metal complexes include iron complexes such as di(triphenylphosphine) iron dichloride, di(tributylamino) iron dichloride, triphenylphosphine iron trichloride, (1-bromo)ethylbenzene-triethoxyphosphine iron dibromide, (1-bromo)ethylbenzene-triphenylphosphine iron dibromide, (1-bromo)ethylbenzene-[4,4'-di(5-nonyl)-2,2'-bipyridine) iron dibromide, (1-bromo)ethylbenzene-tri-n-butylamino iron dibromide, (1-bromo)ethylbenzene-tri-n-butylphosphine iron dibromide, tri-n-butylphosphine iron dibromide, 14.4'-di(5-nonyl)-2.2'-bipyridinel iron dibromide, tetrasikyterrimonium iron(ii) trihalide, dicarponylcyclopentadienyl iron(ii) iodide, dicarponylcyclopentadienyl iron(ii) bromide. dicarbonylcyclopentadienyl iron(II) chloride, dicarbonylindenyl iron(II) iodide, dicarbonylindenyl iron(II) bromide, dicarbonylindenyl iron(ii) chloride, dicarbonylfluorenyl iron(ii) iodide, dicarbonylfluorenyl iron(ii) bromide, dicarbonylfluorenyl iron(II) chloride, 1,3-diisopropyl-4,5-dimethylimidazol-2-ylidene iron chloride, and 1,3-diisopropyl-4,5-dimethylimidazol-2-ylidene iron bromide; ruthenium comptexes such as dicarbonylcyclopentadienyl ruthenium(II) icdide, dicarbonylcyclopentadienyl ruthenium(II) bromide, dicarbonylcyclopentadienyl ruthenium(II) chloride, dicarbonylindenyl ruthenium(II) lodide, dicarbonyfindenyf ruthenium(II) bromide, dicarbonyfindenyf ruthenium(II) chloride, dicarbonyfiluorenvi ruthenium/ii) iodide, dicarbonvifluorenvi ruthenium/ii) bromide, dicarbonvifluorenvi ruthenium(ii) chloride, and dichioro-di-2.6-bisi(dimethytamino)-methyli(u-Na)pyridine ruthenium (II): nickel complexes such as carbonylcyclopentadienyl nickel(II) iodide, carbonylcyclopentadienyl nickel(II) bromide, carbonylcyclopentadienyl nickel(II) chloride, carbonylindenyl nickel(ll) iodide, carbonylindenyl nickel(ll) bromide, carbonylindenyl nickel(ll) chloride, carbonylindenyl nickel(II) iodide, carbonylfluorenyl nickel(II) iodide, carbonylfluorenyl nickel(II) bromide, carbonylfluorenyl nickel(II) chloride, o,o'-di(dimethylaminomethyl)phenyl nickel halide, di-triphenylphosphine nickel dibromide, di(tri-n-butylamino) nickel dibromide, 1,3-diaminophenyl nickel bromide, di(tri-n-butylphosphine) nickel dibromide, and tetra(triphenylphosphine) nickel; molybdenum complexes such as tricarbonyloyclopentadlenyl molybdenum(II) iodide, tricarbonyloyclopenfadieny) molybdenum(ii) bromide, tricarbonylcyclopenfadieny) molybdenum(ii) chloride, di-N-aryl-di(2-dimethylaminomethylphenyl) lithium molybdenum, di-N-aryl(2-dimethylaminomethylphenyl)-methyl lithium molybdenum, di-N-aryl(2-dimethylaminomethylphenyi)-trimethylsilylmethyl lithium molybdenum, and di-N-aryl(2-dimethylaminomethylphenyt)-p-totyl lithium molybdenum; tungsten complexes such as tricarbonylcyclopentadienyl tungsten(II) iodide, tricarbonylcyclopentedienyl tungsten (II) bromide, and tricarbonylcyclopentedienyl tungsten (II) chloride; cobalt complexes such as dicarbonylcyclopentadienyl cobait (i); manganese complexes such as tricarbonylcyclopentadienyl manganese(I) and tricarbonyl(methylcyclopentadienyl) manganese(I); rhenium complexes such as tricarbonylcyclopentadieny! rhenium(I) and dioxobis(triphenylphosphine) rhenium iodide; rhodium complex salts such as tri(triphenylphosphine) rhodium chloride; palladium complexes such as triphenylphosphineacetyl palladium; copper complex salts of diphenanthroline, substituted diphenthroline, 2.2': 6',2"-terpyridine, pyridinimine, or cross-tinked aliphatic diamine, and copper complexes such as acetyl/4-4'-dit5-nonyli-2.2'-bipyridine1 copper, hexafluorophosphine-dii4-4'-di (5-nonvi)-2.2'-bicyridinel copper, copper thiocyanate, bipyridine copper with coordinated O. S. Se, or Te, and copper with coordinated alkylbipyridinylamine, alkyl substituted tripyridine, di(alkylamine)alkylpyridine, iminodipyridine, ethylenediamined/pyridine, or tri(pyridinylmethyl)amine. Of these, dicarbonylcyclopentadienyl iron (I) iodide, dicarbonylcyclopentadieny: ruthenium(II) iodide, and carbonylcyclopentadieny! nickel(II) iodide and the like are preferred. These transition metal complexes can be used either singularly, or in combinations of two or more different complexes.

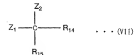
[0042] The organishalogen compound used in the production of a copolymer of the present invention includes compounds comprising from 1 to 4, or even more, halogen atoms (such as fluorine, chlorine, bromine, and lodine). This organishalogen compound functions as an initiator for initiating the polymerization reaction by interacting with the transition metal complex to generate radicals. This type of organishalogen compound can be used either singularly, or in a combination of two or more different compounds. There are no particular restrictions on the organization compound, although suitable examples include the halogen compounds represented by the formulas (VII) and (VIII) shown below.

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$$Z_1 - SO_2 - R_{16}$$
 ... (VIII)

[0043] In these formulas, the groups R₁₄ and R₁₅ may be either the same or different, and each represent a hydrogen atom, alkyl group, cycloalkyl group, aryl group, aralkyl group, or an organic group containing a hetero atom. The group Ris represents an alkyl group, cycloalkyl group, aryl group, aralkyl group, or an organic group containing a hetero atom. The alkyl groups, cycloalkyl groups, aryl groups, aratkyl groups, or organic groups containing a hetero atom represented by R₁₄ to R₁₆ may contain substituent groups. The group Z₁ represents either a halogen atom or an organic group containing a hatogen atom, and the group Z₂ represents either the same as Z₁ or the same as R₁₄ and R₁₈. Examples of suitable halogen atoms include fluorine, chlorine, bromine, and jodine, although chlorine, bromine, and locine are particularly preferred. Specific examples of the above alkyl groups include C1 to C12 alkyl groups such as methyl groups, ethyl groups, n-propyl groups, isopropyl groups, n-butyl groups, isobutyl groups, sec-butyl groups, and t-butyl groups. Specific examples of the excloalkyl groups include C4 to C12 cycloalkyl groups (and preferably C4 to C8 cycloalkyl groups) such as cyclopentyl groups, cyclohexyl groups, and cyclooctyl groups. Specific examples of the aryl groups include C6 to C12 aryl groups such as phenyl groups, tolyl groups, and naphthyl groups, and specific examples of the aralkyl groups include C7 to C14 aralkyl groups such as benzyl groups and phenethyl groups. Examples of the organic groups containing a hetero atom include organic groups containing at least one hetero atom (such as a nitrogen, oxygen, or sulfur atom), and specific examples include ester groups [including alkoxycarbonvi groups (including aliphatic C1 to C10 alkoxycarbonyl groups such as methoxycarbonyl groups, ethoxycarbonyl groups, propoxyearbonyl groups, and butoxyearbonyl groups; and C6 to C12 aryloxyearbonyl groups such as phenoxyearbonyl groups); acytoxy groups (including aliphatic C2 to C10 acytoxy groups such as acetyloxy groups and propionyloxy groups); and C6 to C12 grylcarbonyloxy groups such as benzovioxy groups], ketone groups (including allphatic C1 to C10 acyl groups such as formyl groups and acetyl groups; and C6 to C12 arylcarbonyl groups such as benzoyl groups), ether groups (including aliphatic C1 to C10 alkoxy groups such as methoxy groups and ethoxy groups; and C6 to C12 aryloxy groups such as phenoxy groups and naphthoxy groups), as well as carboxyl groups, hydroxyl groups, amino groups, amide groups, iming groups, cyano groups, nitro groups, thioaster groups, thioketone groups, and thioether groups (such as alky) sulfide groups and anyl sulfide groups). The non-halogen atoms of groups Rt4 and Rtx may contain the same types of substituents allowed in the aforementioned hydrocarbon ligands.

[0044] Examples of preferred halogen compounds include halogenated hydrocarbons represented by the above formula (VII), halogenated esters (halogen-containing esters), halogenated ketones (halogen-containing ketones), and suffray halides (halogenated suffony) compounds) represented by the above formula (VI). Specific examples of halogenated hydrocarbons include haloalkanes (including C1 to C12 alkyl monohalides such as methyl chloride, methyl bromide, embryl bromide, entryl oldide, seprepyl bromide, propyl bromide, bronzyl chiloride, benzylydryl chiloride, bromyl bromide, chilorophoraymena, and dichlorodbromylydryl bromide.

and, Specific examples of halogen-containing esters include C1 to C10 alkyl esters, substituted alkyl esters, or alkenyl esters of halogen-containing C2 to C12 monocarboxylic acids (and preferably C1 to C4 alkyl esters of halogen-containing C4 to C8 monocarboxylic acids) such as methyl dichloroacetate, methyl trichloroacetate, nylacetate, ethyl 2-bromo-2-methylpropionate, hydroxyethyl 2-bromopropionate, glycidylmethyl 2-bromopropionate, properly! 2-bromopropionate, vinyt chloroacetate, bromolactone, p-carboxyphenylethy! 2-bromopropionate, methyl 2-chloroisobutyrate, ethyl 2-chloroisobutyrate, methyl 2-bromoisobutyrate, ethyl 2-bromoisobutyrate, methyl 2-iodoisobutyrate, and ethyl 2-iodoisobutyrate; C1 to C10 alkyl esters of halogen-containing C1 to C14 polycarboxylic acids (and preferably C1 to C4 alkyl esters of halogen-containing C2 to C8 dicarboxylic acids) such as dimethyl 2-chloro-2-methylmalonate, diethyl 2-chioro-2-methylmalonate, dimethyl 2-bromo-2-methylmalonate, diethyl 2-bromo-2-methylmalonate, diethylmalonate, die ylmalonate, dimethyl 2-lodo-2-methylmalonate, diethyl 2-lodo-2-methylmalonate, and dimethyl 2-bromo-2,4.4-trimethyigitatrate; and halogen-containing C2 to C12 carboxylic acids such as dichloroacetic acid, dibromoacetic acid, 2-chioroisobutyric acid, and 2-bromoisobutyric acid. Specific examples of halogen-containing ketones include halogenated C1 to C10 alkyl-C1 to C10 alkyl ketones such as 2-chloroacetone, 1,1-dichloroacetone, ethylchloro methyl ketone, and 1-bromoethyl ethyl ketone; and halogenated C1 to C10 alkyl-C6 to C12 aryl ketones such as 2,2-dichloroacetophenone and 2-bromoisobutyrophenone. Specific examples of halogen-containing netriles include 2-bromopropionitrile, and benzyl thiocyanate, which can also be considered a part of this series. In addition to the halogen compounds represented by the above formula (VII), organohalogen compounds containing 3 or 4 halogen atoms can also be used as reaction initiators. Examples of compounds containing 3 halogen atoms include C1 to C12 trihaloalkanes such as chloroform, C7 to C14 aralkyl trihalides such as trichforophenylmethane, C1 to C10 alkyl esters of halogen-containing C2 to C12 monocarboxylic acids such as acetyltrichloromethane, and halogenated C1 to C10 alkyl-C1 to C10 alkyl ketones such as 1.1,1-trichloroacetone, whereas examples of compounds containing 4 halogen atoms include C1 to C12 tetrahaioalkanes such as carbon tetrachloride and bromotrichloromethane. Even compounds with more than 4 halogen atoms such as trifluorotrichioroethane can be used.

[0045] Specific examples of sulfonyl halides represented by the above formula (VIII) include allighatic sulfonyl chicrides (and particularly alkanesulfonyl halides such as methanesulfonyl halides) such as methanesulfonyl chloride, methanesulfonyl bromide, methanesulfonyl iodide, chloromethanesulfonyl chloride, chloromethanesulfonyl bromide, chloromethanesulfonyl iodide, dichloromethanesulfonyl chloride, dichloromethanesulfonyl bromide, dichloromethanesulfonyl jodice, bromomethanesulfonyl chloride, bromomethanesulfonyl bromide, bromomethanesulfonyl jodide, dibromomethanesulfonyl chloride, dibromomethanesulfonyl bromide, dibromomethanesulfonyl iodide, iodomethanesulfonyl chloride, iodomethanesulfonyl bromide, iodomethanesulfonyl iodide, diiodomethanesulfonyl chloride, diiodomethanesulfonyl bromide, diiodomethanesulfonyl iodide, and trichloromethanesulfonyl chloride; and aromatic sulfonyl halides such as benzenesulfonyl chloride, benzenesulfonyl bromide, benzenesulfonyl iodide, p-methylbenzenesulfonyl chloride, p-methylbenzenesulfonyl bromide, p-methylbenzenesulfonyl iodide, p-chlorobenzenesulfonyl chloride, p-chlorobenzenesulfonyl bromide, p-chlorobenzenesulfonyl lodide, p-methoxybenzenesulfonyl chloride, pmethoxybenzenesulfonyl bromide, p-methoxybenzenesulfonyl lodide, p-nitrobenzenesulfonyl chloride, p-nitrobenzenesulfonyl promide, p-nitrobenzenesulfonyl p-nitrob nesulfonyl bromide, p-nitrobenzenesulfonyl iodide, p-fluorobenzenesulfonyl chloride, p-carboxybenzenesulfonyl chloride, ride, p-aminodiazobenzenesulfonyl chloride, 2,5-dichiorobenzenesulfonyl chloride, 2,5-dimethoxybenzenesulfonyl chloride, 2-hydroxy-3,5-dichlorobenzenesulfonyl chloride, 1-naphthalenesulfonyl chloride, 2-naphthalenesulfonyl chloride, ride, (5-amino-2-naphthalene)sulfonyl chloride, 1,4-disulfonylchloride benzene, 1,4-disulfonyldibromide benzene, 1,4-disulfonyldilodide benzene, 2,6-disulfonyldichloride naphthalene, 2.6-disulfonyldibromide naphthalene, and 2.6-disulfor vidiodide naphthalene. In addition, examples of other halogen compounds that contain hetero atoms include hatogeneted alcohols (including hatogeneted C1 to C10 aliphatic, allovoile, and aromatic alcohols such as 2.2-dichioroethanol and 2,2-dibromoethanol), halogenated nitrites (including halogenated nitrites such as dichloroacetonitrite and dibromoacetonitrile), halogenated aldehydes, and halogenated amides.

[0046] In a living radical polymerization, a Lewis acid and/or an emine, which act as activating agents by acting upon the metal complex to accelerate the radical polymerization, can also be added. Such Lewis acids end amines can be used either singularly, or in combinations of two or more different compounds. There are no particular restrictions on the type of Lewis acid used, and suitable examples include the Lewis acids represented by the formulas (IX) and (X) shown below.

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$$R_{17}$$
 M_2 R_{20} M_2 R_{20} R_{17}

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[0047] In these formulas, M₁ represents an element from either group 3 or group 13 of the periodic table, and M₂ represents an element from either group 4 or from prioridic table. The groups R₁-Lis R₂ may be either the same or different, and each represent a helogen atom, or an alkyl group, cycloalkyl group, anyloxy group, or analkyl group, cycloalkyl group, anyloxy group, or analkyloxy group, specific examples of the aforementioned M₁ include elements from group 3 of the periodic table such as scandium Sc and yttrium Y, and elements from group 13 of the periodic table such as bronn B, aluminum Al, gallium Ga, and indium In, and of these, Sc, B, and Al are preferency and Sc and Al are particularly described. Specific examples of the aforementoned M₂ include elements from group 4 of the periodic table such as stanium T1, zirconium Z1, and hafnium III, and elements from group 14 of the periodic table such as stillown S1, and then S1. Z2, and S1 are prefered.

[0048] In the case of the groups R_{17} to R_{20} , specific examples of the alkyl groups include C1 to C12 alkyl groups such as mathlyl groups, eithyl groups, ethyl groups, septicity, sopropyl groups, septicity, sopropyl groups, specific examples of the eyoloalkyl groups include C4 to C12 cycloalkyl groups (and preferably C4 to C8 cycloalkyl groups) such as cyclopentyl groups, explorely groups, and cyclopentyl groups, such as cyclopentyl groups, and cyclopedyl groups, such as cyclopentyl groups, such as the partial groups include C6 to C12 ay groups such as benzyl groups and phonethyl groups. And specific examples of the arial groups include C7 to C14 aralkyl groups such as benzyl groups and phonethyl groups. Furthermore, specific examples of the cycloalkyloxy groups include C7 to C14 aralkyl groups with as cyclopentyloxy groups. According to examples of the aralkyloxy groups include C7 to C14 aralkyloxy groups such as benzyloxy groups and phonethyloxy groups and proposed to the cycloalkyloxy groups and phonethyloxy groups.

[0049] Those groups R₁₇ to R₂₆ that are not halogen atoms may contain the same types of substituents ellowed in the altermenticed hydrocarbon ligands. For example, anyloxy groups may contain a substituent group such as one or more C1 to C5 alkyl groups on the aromatic ring. Specific examples of such substituted anyloxy groups include 2-methylphenoxy groups, 3-methylphenoxy groups, 4-methylphenoxy groups, 2-de-dihylphenoxy g

[0050] Specific examples of the compounds represented by the above formula (IX) inolude aluminum-based Lowis acids such as aluminum alikoxides [including C1 to C4 alkoxides or aryloxides of aluminum such as aluminum trethoide, aluminum trisopropoxide, aluminum triva-buloxide, aluminum triva-buloxide, and aluminum triphenoxide; and alikylaluminum aryloxides such as methylaluminum bis(2,8-di-l-bulytyhenoxide), ethylaluminum bis(2,8-di-l-bulytyhenoxide), and methylaluminum bis(2,8-di-l-bulytyhenoxide), and methylaluminum bis(2,8-di-bulytyhenoxide), and aluminum trifichide, aluminum triboride, and siluminum triboride, and scandium t

[0051] Specific examples of the compounds represented by the above formula (X) include litanium-based Lowis acids, including thanium alkoxides (such as titanium tetra-methoxide, titanium tetra-methoxide, titanium tetra-propoxide, titanium tetra-hotoxide, titanium tetra-hotoxide, titanium tetra-propoxide, chlorotitanium disopropoxide, and trichiorotitanium sopropoxide); titanium tetra-bioxide, chlorotitanium disopropoxide, and trichiorotitanium isopropoxide); titanium tetra-disordis, chlorotitanium tetra-propoxide, chwis acids titanium tetra-propoxide, zirconium-tetra-to-methoxide, zirconium tetra-propoxide, and zirconium tetra-propoxide and tin-based Lowis acids that correspond with the above titanium-based Lowis acids (including tin alkoxides such as tin tetra-titoride, in tetra-promide, and tin-foreided).

[0052] Of these compounds, preferred Lewis acids include metal compounds (and particularly metal alkoxides) in which the metal is selected from a group consisting of atuminum, scandium, titanium, zirconium, and tin, and specific

examples of proferred compounds include aluminum alkoxidas (such as aluminum triethoxide, aluminum triiscoropoxide, aluminum tris-butoxide, and aluminum tri-t-butoxide), scandium alkoxides (such as scandium triscopropoxide), itämum alkoxides (such as titanium tetramboxide, titanium tetramboxide, blanium tetraiscopropoxide, titanium tetran-butoxide, titanium tetra-t-butoxide, and titanium tetraphenoxide), ziroorium alkoxides (such as ziroonium tetraiscopropoxide), and fin alkoxides (such as in tetraiscopropoxide) and

10053] Furthermore, there are no particular restrictions on the type of amine that can be used and any nitrogencontaining compound can be used, including secondary amines, tertiary amines, and nitrogen-containing amountatheterocyclic compounds, although secondary amines and tertiary amines are preferred. Specific examples of suitable secondary amines include dimethylamine, diethylamine, di-p-propylamine, disopropylamine, di-n-butylamine, giventine, piperidine, 22,66-tertemethylippieridine, piperazine, and morpholine, whereas specific examples of suitable retiral or amines include timethylamine, triethylamine, trien-propylamine, 11-n-butylamine, disopropylethylamine, NN-NV, N-tetramethylethylenediamine, 11-disclarabloyled, 43,0]non-5-en, 14-disabloyled, 22 glorate, and 1,8-disabloyled [54,0]undec-7-ene. Furthermore, compounds containing two or more of a primary amine portion, a secondary amine portion, and a tertiary amine portion within the one molecule can also be used. Specific examples of such compounds include diethylenetiamine, triethylenetiamine, triethylenetiamine, end 44-(2-aminosthylippieridine. The ratio etween the transition metal complex and the Lewis acid or amine is typically within a range from former/laiter = 0.05/1 to 101/finding articlo, and preferably from 0.11/t to 51/firmiar ratio).

[0054] Furthermore, Ilving radical polymerization can also be conducted using stable radicals. Examples of such stable radicals include mixtures of a stable free radical compound and a radical polymerization initiator, or any of the various allowownines

[0055] A stable free radical compound is one that can exist alone as a stable free group at room temperature or under the polymerization conditions, or is able to meat with a growing, non-terminated radical during a polymerization reaction, generating a bond that is capable of re-dissociating. Specific examples include compounds capable of generating one or more intoxide radicals or hydraxyl radicals such as 2.2.6.8-tetramethyl-1-piperidinyloxy (TEMPO), 4-anno-2.2.6.6-tetramethyl-1-piperidinyloxy, 4-dimenthyl-1-piperidinyloxy, 4-dimenthyl-1

[0056] The ratical polymorization initiator can be any compound that undergoes discomposition to generate free raticals and specific examples include are compounds such as 2.2 azoblesioputyronifite [2.2-zaobles[2.4-dimethylvaleronitrite], discyl peroxides such as benzoyl peroxide, ketone peroxides such as methyl ethyl ketone peroxide, peroxyketals such as 1.7-ble(t-butyberoxy) 3.3.5-trimethylcyclohexane, hydroperoxides such as cumene hydroperoxide, d, dallyl peroxides such as dicumyl peroxide, and organic peroxides of peroxyketers such as 1-butyl peroxypivaliate and 1-butyl peroxybenzate. Furthermore, conventional polymertzation accelerators such as dimethylaniline or cobalt nephthenate that are commonly combined with organic peroxides can also be used.

[0057] The radical polymerization initiator is typically used in a quantity equivalent to 0.05 to 5 mols, and preferably from 0.2 to 2 mols, per 1 mol of the aforementioned stable free radical compound.

[0058] Examples of alkoxyamines include those compounds disclosed on page 107 of "Handbook of Radical Polymerization", NTS Co., Lot, and those disclosed in J. Am. Chem. Soc., 1999, 121, 3904, and of these, the compounds shown below are particularly preferred.

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- 25 [0059] Specific examples of methods of producing a copolymer using a living radical polymerization include the following methods:
 - (1) a method in which the monomers are added sequentially, wherein once the conversion ratio of the first monomer has reached 100%, the second monomer is added to complete the polymerization, and his process is then repeated, thus generating a block copolymer.
 - (2) a method in which the second monomer is added, and the polymerization is continued, once the degree of polymerization or the molecular weight has reached a target value, regardless of whether or not the conversion ratio of the first monomer has reached 100%, thus generating a gradient copolymer in which random portions exist between the block chains, and.
 - (3) a method in which once the degree of polymerization or the molecular weight has reached a target value, regardless of whether or not the conversion ratio of the first monomer has reached 100%, the reaction is halted, and the polymer is removed from the system and added to another monomer as a macroinitiator, thus allowing a copolymerization to proceed intermittently, generating a block copolymer.
- (dec) [0060] There are no particular restrictions on the polymerization method, and conventional methods such as bulk polymerization, solution polymerization, suspension polymerization, or emulsion polymerization can be employed, although solution polymerization is used, here are no particular restrictions on the solvent, and suitable solvents include aromatic hydrocarbons (such as benzene, toluene, and xylene), alicybic hydrocarbons (such as cyclobravene), aliphatic hydrocarbons (such as benzene, toluene, and xylene), alicybic hydrocarbons (such as sex), as example, and such as decident and such as example, and proferably from 40 to 150°C, and at either normal prosesure or under pressurized conditions.
 - [0061] A living aninoin polymerization is typically conducted using an alkali metal or an organoalkali metal as the polymerization inhilator, either under vacuum or in an inert gas atmosphere such as nitrogen or argon, in an organic solvent, and at a temperature within a range from -100 to 50°C, and proferably from -100 to -20°C. Examples of suitable alkali metals include lithium, potassium, sodium, and cessium, whereas examples of suitable organoalkali metals include alkylatod, eithylated or anyistad compounds of the above alkali metals. Specific examples include or-bulylithium, see country in the country of the

[0662] Suitable organic ablents include the types of organic solvents typically used in anionic polymerizations, including aromatic hydrocarbons (such as benzene toluene, and kylene), aliphalic hydrocarbons (such as hozane and cotane), alicyclic hydrocarbons (such as sozionere toluene, and kylene), ketones (such as acetone, methyl ethyl ketone, and cyclohaxanone), ethers (such as teitarlyddriuran, dioxane, drijviene glycol dimethyl ether, aniole, and identifyere glycol dimethyl ether), aniole, and identifyere glycol dimethyl ether), aniole, and intervaling hosepharamide. Furthermore, other conventional additives, including the alkali metal salts or alkali earth metal salts of mineral acids such as lithium chloride and the like, can also be used for controlling the coopylmetrization reaction.

[0063] In those cases where a compound containing an active hydrogen atom such as a hydroxyl group or a carboxyl group is used in a living anionic polymerization, the active hydrogen atom is first protected using a conventional protective reaction such as a silylation or a conversion to an acetal or BOC group, the polymerization reaction is conducted, and tene following polymerization, a deprotection reaction is conducted using either actid or alkali.

[0064] Progress of the copolymerization reaction, or confirmation of completion of the reaction, can be ascertained assily using techniques such as gas chromatography, lequid chromatography, get permeation chromatography, membrane asmometry, or NMT. Following completion of the copolymerization reaction, the copolymer can be obtained via typical separation and purification methods such as column purification, or by adding the crude product to water or another poor solvent, and then filterino off and drivin the precipitated polymer component.

3) Polymer Electrolyte of the Present Invention

20 [0065] A polymer electrolyte of the present invention comprises a copolymer described above, and an electrolyte salt. The copolymer can use a mixture of two or more copolymers with different structural units.

[0066] There are no particular restrictions on the electrolyte salt used in the present invention, and an electrolyte comprising the ion identified as the most desirable charge carrier can be used, although electrolytes that display a large dissociation constant within the solid polymer electrolyte produced on curing are preferred. Suitable examples include sikali metal salts, quatemary ammonium salts such as (CH₃),NBF_e, quatemary phosphonium salts such as (CH₃),NBF_e quatemary phosphonium salts such as (CH₃),PBF, explored in the salts such as AgO(QL), or protocine coids such as hydrochoric acid, and fuoroboric acid, and of those, alkeli motal salts, quaternary ammonium salts, quaternary phosphonium salts, and transition motal salts are preferred.

[0067] Specific examples of suitable electrolyte salts include conventional alkali motal salts such as LiCF₂So₂, LiN (CF₂So₂)₂, LiC(CF₃So₂)₂, LiC(CF₃So₂)₂, LiC(CF₃So₂)₃, LiC(CF₃So₂)₃, LiN₂C₂So₃, LiN(C₂F₃So₃)₂, LiP₃C₃So₃, LiN(C₂F₃So₃)₃, LiN₃C₃So₃, LiN(C₂F₃So₃)₄, LiC(C₃So₃)₄, LiCF₃So₃, LiN(C₃So₃)₅, LiCF₃So₃, LiN₃NaF₆, NaCiO₄, NaI, NaBF₄, NaSiO₅, NaSF₆, NGC₃O₃, NaPF₆, NGCiO₄D₃, and these electrolyte salts may be used either singularly, or in mixtures of two or more different salts. Of the above electrolyte salts, lithium satis are perticularly preferred.

(2688) The quantity added of the electrolyte salt is typically within a range from 0.05 to 80 mo%, and preferably from 0.01 to 50 mo%, relative to the quantity of alkylene oxide units within the copolymer. A polymer electrolyte of the present invention can be produced by adding, and mixing (complexing) an electrolyte salt with an afforementioned copolymer. There are no particular restrictions on the method used for this addition and complexing process, and suitable methods include a method in which the copolymer and the electrolyte salt are dissolved in a suitable method in which the copolymer and the electrolyte and the affective salt are missolved as the mixing the process. The process is a mixing the process of the process of

[0069] Modling the aforementioned solid polymer electrolyte into sheet, membrane, or film form is particularly desirable, and by so doing, the degree of freedom of the processed surface increases, which provides significant practical benefills. The sheet-like solid polymer electrolyte can be produced by any of a variety of coating techniques inclining roll coating, curtain coating, spin coating, dipping, or casting, and using one of these techniques, a film of the solid polymer electrolyte is formed on the surface of a substrate, and the substrate is subsequently removed to yield the solid polymer electrolyte sheet.

[0070] A solid polymer electrolyte of the present invertion contains a copplymer comprising block chains A. B, and C. In the composition, the block chain A, because it contains a repeating unit represented by the formula (i), can be considered as the polymer segment (P1) that displays ionic conductivity, the block chain B. because it contains a repeating unit represented by the formula (ii), can be considered as a polymer segment (P2) that displays no ionic conductivity, and the block chain C of a preferred embodiment, which contains a repeating unit represented by the formula (iii), is also a polymer segment (P2) that displays no ionic conductivity, and in those cases where the block chains are represented by each of these polymer segments, the copolymer is a polymer in which the segments are arranged in a P2, P1, P2 sequence.

[0071] A solid polymer electrolyte of the present invention functions as an ion conductive firm, and this film comprises a polymer containing a polymer segment (P1) that displays ionic conductivity, and a polymer segment (P2) that displays no ionic conductivity, wherein the film comprises P1 microdomalins and P2 microdomains that form a microphase sep-

arated structure.

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[0072] Furthermore, this microphase separated structure within the ion conductive film is preferably a network-type microphase separated structure. By ensuring such a structure, the ionic conductivity, the physical characteristics, the thermal characteristics, and cancilularly the film strength, can all be improved.

BRIFF DESCRIPTION OF THE DRAWINGS.

[0073] FIG. 1 is a transmission electron microscope photograph showing a cross-section of an electrolyte film comprising a B-A-B type multi-branched polymer compound according to the present invention.

BEST MODE FOR CARRYING OUT THE INVENTION

[0074] As follows is a more detailed description of the present invention using a series of examples, although the present invention is in no way restricted to the examples presented below.

Example 1 <Solid polymer electrolyte comprising a B-A-B type multi-branched polymer compound in which the block chain A is poly-methoxypolyethylene glycol monomethacrylate, and the block chain B is polystyrene>

(1) Synthesis of the block chain A

[0075] To 89.40 g of totuene under an argon atmosphere were added 0.65 g (0.05 mmol) of dichlororis(triphenylphosphine) ruthenium and 22.35 g (20.0 mmol) of methoxypolyethylene gycoid monomethacrylate (Blemer PME-1000, a compound of the aforementioned formula (IV) wherein m = 23, manufactured by NOF Corporation), and following mixing to generate a homogeneous solution, 0.03 g (0.2 mmol) of di-n-bufylamine and 0.02 g (0.1 mmol) of 2.2-dichloroacetophenone were added. The resulting mixture was then heated to 80°C under constant stirring to initiate the
polymetization reaction. Following reaction for 22 hours from the point of reaction initiation, the polymerization reaction was halted by cooling the polymerization reaction system to 0°C. The polymerization ratio was 55.0%. The polymerization solution was then purified by passage through a column to remove the metal complex and any unreacted monomers, and the following was then removed under reduced pressure, yielding poly-methoxypolyethylene glycol monomethacrylate (hereafter abbreviated as P-PME1000-1). The resulting P-PME1000-1 was a single peak polymer with a number average molecular weight (MP) of 125.500.

(2) Synthesis of a B-A-B type multi-branched polymer compound

(28 [0078] To 20.37 g of foluene under an argon atmosphere were added 0.01 g (0.01 mmol) of chloropeniamethylogoclopentadiaderyi-bis(triphenylphosphine)ruthenium, 6.13 g (0.05 mmol) of P-PME1000-1, 2.60 g (2.5 mmol) of syvene, and 0.57 g (5.0 mmol) of n-octane, and following mixing to generate a homogeneous solution, 0.01 g (0.1 mmol) of dinebutylamine was added, and the resulting mixture was heated to 100°C under constant stirring to initiate the polymerization reaction reaction in greation for 20 bours from the point of reaction initiation, the polymerization reaction system to 0°C. The styrene polymerization ratio was 26%. The polymerization value to purified by passage through a column to remove the metal complex and any unreaded monomers, and the toluene was then removed under reduced pressure, yielding a multi-branched polymer compound with a poly-(styreno-b-PME-1000-b-styreno) by postmouture in which the methoxypolyethylene glycol acted as a graft chain. The thus obtained multi-branched polymer compound was a single peak polymer with a ratio between the block chain 8 and the block chain 8 of AB = 2.067 (degree of polymerization ratio), and Mn = 135,000. Furthermore, the thus obtained multi-branched polymer compound was dissolved in accitors and used to generate a film, and inspection of the cross-section of this film under a transmission electron microscope (TEM) revealed a network-type microphase separated structure.

50 (3) Preparation, and evaluation of the properties of a solid polymer electrolyte film

[0077] I g of the multi-branched polymer compound obtained by the process described above was dissolved in 25 ml of action under an argon atmosphere, and 0.09 g of LLGO, as an electrolyte salt was then added to the resulting solution and dissolved to generate a homogeneous solution. The solution was then spread onto a Tellon (registered trademark) substrate, allowed to stand for 24 hours at room temperature, and was then dired under reduced pressure at 80°C for 24 hours, thus generating a uniform solid electrolyte film (film thickness 140 µm). Under an argon atmosphere, this film was then sandwiched between platinum plates and an impedance analyzer of frequency 5 Hz to 10 MHz (Solarton-1200) was used to measure the ionic conductivity by complex incedance analysis. The measurement

revealed an ionic conductivity at 23°C of 3.8 × 10.4 S/cm.

[0078] Furthermore, the cross-section of the thus obtained solid polymer electrolyte was also inspected using a transmission electron microscope (TEM). The resulting photograph is shown in FIG. 1. From FIG. 1 it is evident that the structure within the lim is a network-two microphase securated structure.

Example 2 <Solid polymer electrolyte comprising a S-A-B type multi-branched polymer compound in which the block chain A is poly-methoxypolyethytene glycol monomethacrylate, and the block chain B is polystyrene>

(1) Synthesis of the block chain A

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[0079] To 115.80 g of toluene under an argon atmosphere were added 0.10 g (0.1 mmol) of dichlorotris(triphenylphosphine)uthenium and 28.5 g (26.0 mmol) of Blemer PME-1000, and following mixing to generate a homogeneous
solution. 0.05 g (0.4 mmol) of d-n-bulylamine and 0.04 g (0.2 mmol) of 2,2-dichloroxectophoneno were added. The
resulting mixture was then heated to 80°C under constant stirring to initiate the polymerization reaction. Following
reaction for 48 hours from the point of treaction initiation, the polymerization was halted by pooling the polymerization reaction was halted by pooling the polymerization reaction was halted by pooling the polymerization treaction system to 0°C. The polymerization ratio was 50%. The polymerization solution was then curlified by
passage through a column to remove the metal complex and any unreacted monoments, and the tolurine was then
removed under reduced pressure, yielding poly-methoxypolyethylene glycol monomethacyleid (hereafter abbreviated
as P-PME1000-2). The resulting P-PME1000-2 was a single peak polymer with a number average molocular weight
(Mh) of 70.000 (d degree of polymerization of 63).

(2) Synthesis of a B-A-B type multi-branched polymer compound

[0060] To 32.62 g of toluene under an argon atmosphere were added 0.02 g (0.02 mmol) of chloropentamethyloropentadens-bishtiphenylothephne) urbenium, 7.00 g (0.10 mmol) of p-Mtc1000-2, 6.88 g (6.7 mmol) of stylene, and 0.57 g (5.0 mmol) of n-octane, and following mixing to generate a homogeneous solution, 0.03 g (0.2 mmol) of di-houly lamine was added, and the resulting mixinum was heated to 100°C under constant stirring to initiate the polymerization reaction for 45 hours from the point of reaction initiation, the polymerization reaction was hailed by cocling the polymerization reaction system to 0°C. The styrene polymerization conversion ratio was 57%. The polymerization solution was then purified by passage through a column to remove the metal complex and any unreacted monomers, and the solvent was then removed under reduced pressure, yielding a multi-branched polymer compound with a poly-telyrene-bMt-1000-5 telyrene) type structure in which the methoxypolyethylene glycol code as a graft chain. The thus obtained multi-branched polymer compound was a single peak polymer with a ratio between the block chain A and the block chain B of V/RB = 16 (degree of polymerization and the 1000 column).

(3) Preparation, and evaluation of the properties of a solid polymer electrolyte film

[0081] With the exception of using 0.07 g of LiCiO₄, a uniform solid electrolyte film was prepared in the same manner as the example 1, and the ionic conductivity was measured. The measurement revealed an ionic conductivity at 23°C of 5 < 10°3 S/cm. Furthermore, inspection of the thus prepared film using a transmission electron microscope revealed a similar network-type microphase separated structure to that of the solid polymer electrolyte obtained in the example 1.

Example 3 <Solid polymer electrolyte comprising a B-A-B type multi-branched polymer compound in which the block chain A is poly-methoxypolyethylene glycol monomethacrylate, and the block chain B is polystyrene>

(1) Synthesis of the block chain A

[0082] To 69.53 g of I toluren under an argen atmosphere were added 0.05 g (0.5 mmpl) of capper (I) chlorica, 0.16 g (1.0 mmpl) of blyridine, and 29.80 g (60 mmol) of methoxypolyethylene glycol monomethacrylate (Blemer PME-400, a compound of the aforementioned formula (V) wherein m = 9, manufactured by NOF Corporation), and following mixing to generate a homogeneous solution, 0.09 g (0.5 mmol) of 2.2-dichloreacetophenone was added, and the resulting mixture was then heated to 80°C under constant stirring to initiate the polymerization reaction reaction reaction in a point of reaction initiation, the polymerization reaction was haited by cooling the polymerization reaction system to 0°C. The polymerization conversion ratio was 90%, 500 ml of THF was then added to the reaction solution, excess magnesium sultate was added, and following filtering, the filterite was purified by passage through a column to remove the metal complex and any unreacted monomers. The solvent was then removed under reduced pressure, yielding poly-methoxypolyethylene glycol monomethacrylate (hereafter abbreviated as P-PME400). The resulting P-PME400 was a single peak polymer with Min = 50.000 (a degree of ophyrmetrization of 110).

(2) Synthesis of a B-A-B type multi-branched polymer compound

1083 To 14.58 g of tolurene under an argon atmosphere were added 0.01 g (0.1 mmol) of copper (I) chloride, 0.03 g (0.2 mmol) of bipyridine, 5.00 g (0.9 mmol) of P-PME400, 1.25 g (12 mmol) of styrene, and 0.11 g (1.0 mmol) of nociare, and following mixing to generate a homogeneous solution, the resulting mixture was heated to 100°C under constant stirring to initiate the polymerization reaction. Following reaction for 25 hours from the point of traction installation, the polymerization reaction was halted by cooling the polymerization reaction system to 0°C. The styrene polymerization conversion matio was 70%. The reaction solution was then purified by passage through a column for remove the metal complex, and the solvent was then removed under reduced pressure, yielding a multi-branched polymer compound with a poly-(styrene-b-PME-400-b-styrene) type structure in which the methoxypolyethylene glycol acted as a graft chain. The thus obtained multi-branched polymer compound was a single peak polymer with a ratio between the block chain 8 of 4/8 = 12/1 (degree of colymerization reaction ratio), and Mn = 65.000.

(3) Preparation, and evaluation of the properties of a solid polymer electrolyte film

[0084] With the exception of using 0.08 g of LICIO₄, the ionic conductivity was measured in the same manner as the example 1. The measurement revealed an ionic conductivity at 23°C of 1 × 10⁻⁴ S/cm. Furthermore, inspection of the thus prepared film using a transmission electron microscope revealed a similar network-type microphase separated structure to that of the solid polymer electrolyte obtained in the example 1.

Example 4 <Solid polymer electrolyte comprising a B-A-C type multi-branched polymer compound in which the block chain A is poly-methoxypolyethylene glycol monoacrylate, the block chain B is polystyrene, and the block chain C is isobornyl acrylate>

(1) Synthesis of the block chain B

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[0085] To 48 g of toluene that had already undergone argon degassing treatment were added, under an argon almosphere. 0.02 g (0.02 mmol) of clindropentametrylcycloperitadienyl-bis(triphenylcybasphine)rutherium, 20.8 g (2.00 mmol) of sincepens and 0.57 g (6 mmol) of nectane, and following mixing to generate a homogeneous solution. 300 g (0.4 mmol) of aluminum isopropoxide and 0.08 g (0.4 mmol) of 2-brome isobutyrate were added and mixed to form a homogeneous solution, and the resulting solution was heated to 100°C under constant attring to initiate the polymerization reaction. Following reaction for 50 hours from the point of reaction initiation, the polymerization reaction was halted by cooling the polymerization reaction system to 0°C. The styrene polymerization conversion ratio was 50%. The polymerization solution was then purified by passage through a column to remove the metic complex and any unreacted monomers, and the volatile fraction was then removed under reduced pressure, yielding polystyrene. The trus obtained polystyrene was a sindle peak polymer with Mn ~ 25,000.

(2) Synthesis of a B-A type multi-branched polymer compound

60 [0088] To 88 g of toluene under an argon atmosphere were added 0.02 g (0.02 mmol) of chloropentamethylcyclopentadienyl-bia(ciphenylphosphine)ruthentum, 2.5 g (0.1 mmol) of the polystyrene octained in (1) above, and 271 g (55 mmol) or Inethoxypolyethylare glycol monoacrylate (Blemer AME-400, a compound of the aforementioned formula (VI) wherein m = 9, manufactured by NOF Corporation), and following mixing to generate a homogeneous solution, 0.03 g (0.2 mmol) of clin-britysmine was added, and the resulting mixture was heated to 80°C to initiate the copyoise meization reaction. Following reaction for 55 hours from the point of copolymerization initiation, the copolymerization reaction was shelted by cooling the polymerization reaction system to 0°C. The AME-400 polymerization corporation ratio was 60%. The polymerization solution was then purified by passage through a column to remove the metal complex and any unreacted monomers, and the volatile fraction was then removed under reduced pressure, yielding a multi-branched polymer compound with a poly-cityeneo-b-AME-400) by the structure or which the method through the polymer compound with a poly-cityeneo-b-AME-400) polymer compound was a single peak polymer with Mn = 178.000.

(3) Synthesis of a B-A-C type multi-branched polymer compound

[0087] To 47 g of toluene under an argon atmosphere were added 0.02 g (0.02 mmol) of chloropentamethylcyclopentadionyl-bis(triphenylphosphine)juthenium. 17.8 g (0.1 mmol) of the poly-(styrene-b-AME-400) polymer obtained in (2) above, 3.7 g (18 mmol) of insobornyl acrylate, and 0.57 g (5 mmol) of n-octane, and following mixing to generate a homogeneous solution, 0.03 g (0.2 mmol) of dir-butylamine was added, and the resulting mixture was heated to 30°C.

under constant stirring to initiate the copolymerization reaction. Following reaction for 20 hours from the point of reaction initiation, the copolymentzation reaction was halled by cooling the polymerization reaction system to 0°C. The methyl acrylate conversion ratio was 30%. The polymerization solution was then purified by passage through a column to remove the metal complex and any unreacted monomers, and the volatile fraction was then removed under reduced pressure, yielding a multi-branched polymer compound with a poly-detyrene-hAME-400-b-shoornyal acrylate) type structure in which the methoxypolyethylene glycol acted as a graft chain. The thus obtained multi-branched polymer compound was a singe peak polymer with a ratio between the block chain A, and the combination of the block chain B and the block chair C affects of the properties of t

(4) Preparation, and evaluation of the properties of a solid polymer electrolyte film

[0088] With the exception of using 0.08 g of LiClO₄, a uniform solid electrolyte film was prepared in the same manner as the example 1, and the lonic conductivity was measured. The measurement revealed an ionic conductivity at 23°C of 2 × 10-4 S/cm. Furthermore, inspection of the thus prepared film using a TEM revealed a similar network-type microphase separated structure to that of the solid polymer electrolyte obtained in the example 1.

Example 5

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[0089] Using the B-A-B type multi-branched polymer compound prepared in the example 1, LiClQ, was added in a variety of different molar ratios, as shown in Table 1 (in the Li salt column), relative to each mol of divijene oxide units, and solid polymer electrolytes were prepared in the same manner as the example 1. The conductivity (inel conductivity) of each solid polymer electrolyte was them measured at a number of different temperatures, using the method described in the example 1. The results are shown in Table 1.

Table t

Li salt	Conductivity (S/cm)					
	10°C	20°C	30°C	40°C	50°C	60°C
0.03	7.2E-05	2.0E-04	3.2E-04	4.4E-04	5.4E-04	7.0E-04
0.05	9.2E-05	3.0E-04	4.6E-04	5.6E-04	6.3E-04	8.5E-04
80.0	2.1E-05	8.0E-05	1.8E-04	3.0E-04	4.8E-04	5.5E-04

Comparative Example 1 < Solid polymer electrolyte comprising a A-B-A type multi-branched polymer compound in which the block chain A is poly-methoxypolyethylene glycol monomethacrylate and the block chain B is polystyrene>

[0090] Under conditions including constant stirring under an argon atmosphere at a temperature of -78°C, 2,05 g (19.2 mmol) of styrene was added to 200 ml of a tetrahydrofuran solution containing 0.15 mmol of α-methylstyrene tetramer disodium, and the polymerization reaction was allowed to proceed for 1 hour. When a small sample of the polymerization solution was removed from the reaction system and analyzed by gas chromatography it was evident that the polymerization had finished, and GPC analysis revealed that the product polymer was a single peak polymer with Mn = 13,000. Subsequently, the temperature of the reaction system was raised to -40°C, 0.31 mmol of diphenylethylene was added, and 18.10 g (36.5 mmol) of PME-400 was added droowise to the reaction system. After allowing the copolymerization reaction to proceed for 2 hours, methanol was added to halt the reaction. When a small sample of the polymerization solution was removed from the reaction system and analyzed by gas chromatography it was evident that the polymerization had finished, and GPC analysis revealed that the product copolymer was a single peak polymer with Mn = 130,000. The tetrahydrofuran was then removed from the reaction solution under reduced cressure. yielding a multi-branched polymer compound with a poly-(PME400-b-styrene-b-PME-400) type structure in which the methoxypolyethylene glycol acted as a graft chain. Although this multi-branched polymer compound has substantially the same composition and molecular weight as the multi-branched polymer compound produced in the example 1. because the styrene segment exhibits no pseudo cross-linking effect, the polymer compound was a viscous liquid at temperatures near room temperature, meaning film formation and ionic conductivity measurements were impossible.

Comparative Example 2 < Solid polymer electrolyte comprising a B-A-B type multi-branched polymer compound in which the block chain A is poly-methoxypolyethylane glycol monomethacrylate, and the block chain B is poly-methyl methacrylate.

[0091] To 32.69 g of taluene under an argon atmosphere were added 0.05 g (0.05 mmol) of dichtorrist(triphonyl-phosphins)urbenium, 70 g (0.10 mmol) of methyl method phosphins)urbenium, 70 g (0.10 mmol) of the "PME 1000 obtained in the example 2, 7.01 g (7.0 mmol) of methyl-method phosphins) and 10.57 g (5.0 mmol) of n-octane, and following mixing to generate a homogeneous solution, 0.03 g (0.2 mmol) of din-butylamine was added, and the resulting mixture was heated to 80°C under constant stirring to initiale the polymerization reaction. Following reaction for 20 hours from the point of reaction initiation, the polymerization reaction was helted by cooling the polymerization reaction system to 0°C. The MMA polymerization ratio was 55%. The polymerization solution was then purified by passage through a column to return the metal complex and any unreacted monomers, and the solvent was then removed under reduced pressure, yielding a multi-branched polymer compound with a poly-(MMA-b-PME) to 900-4 MMA be 1000-5 MMA) by set structure in which the methody-vielding in the purphine by the structure in which the methody-vielding in the purphine by the structure in which the methody-vielding in the purphine by the structure in which the methody-vielding in the purphine by the structure in which the methody-vielding in the purphine by the structure in which the methody-vielding in the purphine by the structure in which the method is a significant to the polymer compound was a single peak polymer with a ratio between the block chain a and the block chain 8 of AVB = 1/8 (degree of polymerization ratio), and Mn = 107,000. Furthermore, the this obtained multi-branched polymer compound was dissolved in acetion and used to generate a film, although inspection of the cross-section of this film under a transmission electron microscope revealed a uniform structure, with no evidence of a method structure, with no evi

[0092] Measurement of the ionic conductivity in the same manner as the example 1 revealed a result at 23°C of 2 × 10°6 S/cm.

INDUSTRIAL APPLICABILITY

[0093] A solid polymer electrolyte of the present invention, which exhibits a network-type microphase separated structure as a result of the positioning of segments such as styrene with no ionic conductivity adjacent to ion conductive segments such as polyethylene glycol mono(moth)acrylate, displays practical levels of thermal characteristics, physical characteristics, and ionic conductivity, meaning it has potential uses as the solid polymer electrolyte used in electrochemical elements such as lithium batteries, and bocause the solid polymer electrolyte of the present invention enables reductions in both the size and weight of such elements, it is expected to offer considerable industrial benefits.

Claims

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 A solid polymer electrolyte comprising an electrolyte salt, and a copolymer in which a block chain A containing a repeating unit represented by a formula (I) shown below:

[whorein, R_1 to R_2 each represent, independently, a hydrogen atom or a hydrocarbon group of C1 to C10, R_1 , and R_2 may be bonded logether to form a ring, R_{10} and R_{10} each represent, independently, a hydrogen atom or a methy group, R_3 represents a hydrogen atom or a methy group, R_3 represents a hydrogen atom or group, an acy group, or a silyl group, in represents an integer from 2 to 100, and individual R_{10} and R_{10} groups are either identical or different), a block chain B containing a repealing unit represented by a formula (II) shown below:

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(wherein, R₆ to R₈ each represent, independently, a hydrogen atom or a hydrocarbon group of C1 to C10, and R₉ represents an anyl group), and a block chain C are arranged in a sequence B, A, C.

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A solid polymer electrolyte according to claim 1, wherein said block chain C contains a repeating unit represented by a formula (III) shown below:

$$\begin{array}{c|c}
 & R_{10} & R_{12} \\
 & C & C & C \\
 & R_{11} & R_{12} & \cdots & \cdots \\
\end{array}$$

(wherein, R₁₀ to R₁₂ each represent, independently, a hydrogen atom or a hydrocarbon group of C1 to C10, and R₁₃ represents an aryl group or a heteroaryl group).

- A solid polymer electrolyte according to either one of claim 1 and claim 2, wherein said block chains A to C form a copolymer with a B-A-C bonding sequence.
- A solid polymer electrolyte according to any one of claim 1 through claim 3, wherein a degree of polymerization
 of a repeating unit represented by said formula (f) is at least 10.
 - A solid polymer electrolyte according to any one of claim 1 through claim 4, wherein a degree of polymerization
 of a repeating unit represented by said formula (II) is at least 5.
 - A solid polymer electrolyte according to any one of claim 2 through claim 5, wherein a degree of polymerization of a repeating unit represented by said formula (III) is at least 5.
- A solid polymer electrolyte according to any one of the claim 1 through claim 6, wherein a value of m in said formula
 (i) is an integer from 5 to 100.
 - A solid polymer electrolyte according to any one of claim 1 through claim 7, wherein a value of m in said formula (1) is an integer from 10 to 100.
- A solid polymer electrolyte according to any one of claim 2 through claim 8, wherein said group R₁₃ in said formula (III) is an aryl group, and a degree of polymerization of a repeating unit represented by said formula (III) is at least 5.
 - 10. A solid polymer electrolyte according to any one of claim 1 through claim 9, wherein a molar ratio ((i)/(ii)+C)) between repeating units represented by said formula (i), and a combined total of repeating units represented by said formula (ii) and re-peating units within said block chain C is within a range from 1/30 to 50/1.
 - 11. A solid polymer electrolyte according to any one of claim 2 through claim 9, wherein a molar ratio ((i)'((li)+(ill))) between repealing units represented by said formula (ii), and a combined total of repealing units represented by said formula (iii) are injurially units are injurially
 - A solid polymer electrolyte according to any one of claim 1 through claim 11, wherein a number average molecular weight of said copolymer is within a range from 5,000 to 1,000,000.

- A solid polymer electroly/e according to any one of claim 1 through claim 12, which exhibits a microphase separated structure.
- 14. A solid polymer electrolyte according to any one of claim 1 through claim 13, wherein said electrolyte sall is one or more materials selected from a group consisting of alkali motal, asilts, quaternary ammonium salts, causticinary phosphonium salts, transition metal salts, and protonic acids.

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- 15. A solid polymer electrolyte according to any one of claim 1 through claim 13, wherein said electrolyte sailt is a lithium sailt.
- 16. A copolymer in which a block chain A containing a repeating unit represented by a formula (I) shown below:

$$\begin{array}{c|c} \begin{pmatrix} R_1 & R_3 \\ C & C \end{pmatrix} & \cdots & (I) \\ R_2 & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & \\ & & \\ &$$

(wherein, R_1 to R_2 each represent, independently, a hydrogen atom or a hydrocarbon group of C1 to C10, R_1 and R_2 may be bonded together to form a ring, R_{4a} and R_{1b} each represent, independently, a hydrogen atom or a methy; group, R_2 represents a hydrogen atom, a hydrocarbon group, an acyl group, or a silyl group, or represents an integer from 2 to 100, and individual R_{3a} and R_{3b} groups are either identical or different), a block chain B containing a repealing unit represented by a formula (II) shown below:

(wherein, R₆ to R₈ each represent, independently, a hydrogen atom or a hydrocarbon group of C1 to C10, and R₉ represents an anyl group), and a block chain C containing a repeating unit represented by a formula (III) shown below.

(wherein, $\rm R_{10}$ to $\rm R_{12}$ each represent, independently, a hydrogen atom or a hydrocarbon group of C1 to C10, and $\rm R_{13}$ tepresents an aryl group or a heteroaryl group) are arranged in a sequence B, A, C.

- 55 17. A copolymer according to claim 16, wherein said block chains A to C are bonded together in a B-A-C sequence.
 - A copolymer according to either one of claim 16 and claim 17, wherein a degree of polymerization of a repeating unit represented by said formula (I) is at least 10

- A copolymer according to any one of claim 16 through claim 18, wherein a degree of polymerization of a repeating unit represented by said formula (II) is at least 5.
- A copolymer according to any one of claim 16 through claim 19, wherein a degree of polymerization of a repeating unit represented by said formula (III) is at least 5

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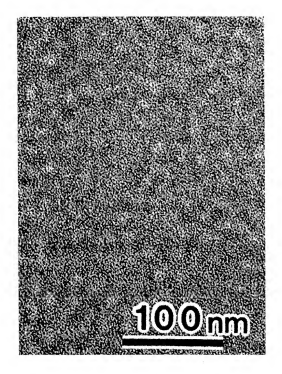
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- A copolymer according to any one of claim 16 through claim 20, wherein a value of m in said formula (I) is an integer from 5 to 100.
- A copolymer according to any one of claim 16 through claim 20, wherein a value of m in said formula (i) is an integer from 10 to 100.
 - 23. A copolymer according to any one of claim 15 through claim 22, wherein said group R₁₈ in said formula (III) is an anyl group, and a degree of polymerization of a repeating unit represented by said formula (III) is at least 5.
 - 24. A copolymer according to any one of claim 16 through claim 23, wherein a motar ratio {(!)/f(i)-f(ii))} between repeating units represented by said formula (i), and a combined total of repeating units represented by said formula (ii) and repeating units represented by said formula (iii) a within a range from 1,30 to 30/1.
- 20 25. A copolymer according to any one of claim 16 through claim 24, wherein a number average molecular weight of said copolymer is within a range from 5,000 to 1,000,000.
 - 26. A copolymer according to any one of claim 16 through claim 25, which exhibits a microphase separated structure.
- 27. A method of producing a copolymer according to any one of claim 15 through claim 26, which utilizes a living radical polymerization in which a transition metal complex is used as a catalyst, and an organohalogen compound comprising 1 or more halogen atoms is used as an initiator.
- 28. An ion conductive film containing a polymer comprising a polymer segment (P1) that displays ionic conductivity and a polymer segment (P2) that displays no ionic conductivity, wherein said film exhibits a notwork-type microphase separated structure.
 - 29. An ion conductive film according to claim 28, wherein said polymer comprising P1 and P2 is a polymer in which said segments are arranged in a P2, P1, P2 sequence.

FIG.1



INTERNATIONAL SEARCH REPORT

International application No. PCT/JP03/09328

CLASSIFICATION OF SUBJECT MATTER Int.Cl' COBF297/00, H01B1/06

According to international Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)
Int.Cl⁷ C08F297/00, H0181/06

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data inseconsulted during the international search (name of data base and, where practicable, search terms used)

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Cimtion of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	JP 10-208545 A (Shin-Etsu Chemical Co., Ltd.),	28,29
a	07 August, 1998 (07.08.98),	1-27
	Claims	
	& US 6096234 A1	
х	JP 11-43523 A (Shin-Etsu Chemical Co., Ltd.),	28,29
A	16 February, 1999 (16.02.99),	1-27
	Claims	
	(Family: none)	
×	JP 10-237143 A (Shin-Etsu Chemical Co., Ltd.),	28,29
A	08 September, 1998 (08.09.98),	1-27
	Claims	
	(Family: none)	
- 1		

15	₹.	Further documents are	listed in the continuation	of Box C.	See	patent famil	y anne
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Date	of the actual completion of the international scarch	Date	of mailing of the international scarcis report

02 September, 2003 (02.09.03) 16 September, 2003 (16.09.03) Name and availing address of the ISA/ Authorized officer Japanese Patent Office

Telephone No.

Form PCT/ISA/210 (second sheet) (July 1998)

Facsimile No.

INTERNATIONAL SEARCH REPORT

International application No. PCT/JF03/09328

(Continua	ion). DOCUMENTS CONSIDERED TO BE RELEVANT	
ategory*	Citation of document, with indication, where appropriate, of the relevant passages	s Relevant to claim N
X	CP 3-19540? A (Upe Industries, Etd.), Z7 August, 1991 (27.08.91), Claims (Family: none)	29,29 1-27
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